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Single-molecule compaction of megabase-long chromatin molecules by multivalent cations

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ABSTRACT

To gain insight into the conformational properties and compaction of megabase-long chromatin molecules, we reconstituted chromatin from T4 phage DNA (165 kb) and recombinant human histone octamers (HO). The unimolecular compaction, induced by divalent Mg2+ or tetravalent spermine 4+ cations, studied by single-molecule fluorescence microscopy (FM) and dynamic light scattering (DLS) techniques, resulted in the formation of 250–400 nm chromatin condensates. The compaction on this scale of DNA size is comparable to that of chromatin topologically associated domains (TAD) in vivo. Variation of HO loading revealed a number of unique features related to the efficiency of chromatin compaction by multivalent cations, the mechanism of compaction, and the character of partly compact chromatin structures. The observations may be relevant for how DNA accessibility in chromatin is maintained. Compaction of saturated chromatin, in turn, is accompanied by an intrachain segregation at the level of single chromatin molecules, suggesting an intriguing scenario of selective activation/deactivation of DNA as a result of chromatin fiber heterogeneity due to the nucleosome positioning. We suggest that this chromatin, reconstituted on megabase-long DNA because of its large size, is a useful model of eukaryotic chromatin.

INTRODUCTION

DNA compaction within the confined space of the eukaryotic cell nucleus in the form of chromosomes is a hierarchical process of outstanding biological significance for storage of genomic DNA molecules, which are several hundreds of megabases long. A well-defined nucleosome core particle (NCP) is formed by wrapping of about 147 bp DNA in 1.75 turns around the histone octamer (HO) (Figure 1A) (1,2). The HO consists of two copies each of the core histones H2A, H2B, H3 and H4. Figure 1A illustrates that the NCP is a polyanion-polycation complex with a net negative charge of about $-146e$, resulting from the negatively charged central particle (with $-294e$ from DNA and $+52e$ from the globular part of the HO). Protruding out from the HO core are ten flexible and positively charged histone tails with a net charge $+96e$, eight of which are N-terminal tails and two C-terminal H2A tails. The NCPs are wedge shaped particles of 11 nm diameter and 6 nm height. The NCPs connected by linker DNA of variable length (20–70 bp) in extended conformation are known as ‘beads on a string’ 10 nm fiber, where the nucleosomes are the fundamental repeating units of chromatin. Nucleosome formation results in diminishing of the effective DNA length. The packing of the 10 nm fiber requires charge neutralization and internucleosomal contacts. Nucleosomes represent the elementary functional assembly for selective access to the DNA genetic information via compaction-decompaction that is mediated by a variety of proteins including linker histones, polyelectrolyte behavior in the presence of multivalent cations that induce condensation and re-dissolution (5,6). Most of the

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studies examined condensation/aggregation of chromatin predominantly at the level of nucleosomes or short nucleosome arrays. Particularly, \textit{in vitro} reconstituted nucleosome arrays using DNA array templates based on the Widom ‘601’ high affinity nucleosome positioning sequence, and with less than a hundred HOs bound per DNA molecule were successfully studied to address the structure of the 30 nm chromatin fiber, chromatin compaction and aggregation (7–9). Other studies particularly focused on the important role of structural recognition of certain DNA sequences by histone proteins, their positioning, and gene regulation (10).

Studies of eukaryotic genomes by chromatin conformation capture-based methods (11) and \textit{in situ} fluorescence techniques (12) suggest that interphase chromatin is organized into cell-type specific chromosome territories (13), occupied by individual chromosomes. Each chromosome territory consists of independent and dynamic topologically associated domains (TAD) (14,15). In human nuclei, TADs are around one megabase-long chromosomal fragments of around 300 or 240 nm in radius, representing euchromatin or heterochromatin regions, respectively (16). At present, studies of long DNA molecules reconstituted with several hundred HOs forming megabase-long chromatin are scant (17,18), and further \textit{in vitro} physico-chemical studies are important to improve our understanding of chromatin organization in nuclei of living cells.

\textit{In vitro} studies on model systems showed that the DNA compaction mechanism and the accompanied changes of DNA biological activity differ dramatically between short and long (>10 kbp) DNAs (19). For example, long-chain DNA exhibits a first order phase transition at the level of individual molecules upon addition of multivalent cations (20), while the compaction of short DNA molecules exhibits a gradual aggregation of DNA into a water-insoluble complex. Consequently, the discrete conformational transition of long DNA abruptly ‘switches off’ the DNA transcriptional activity at the point of the compaction, whereas the genetic activity of short DNA correlates with the degree of DNA aggregation (21). To gain a deeper insight into the problem of long DNA chain compaction in the presence of nucleosome-forming HOs, studies on DNA compaction and transcription by histone-mimicking cationic dendrimers (22) and cationic nanoparticles of several sizes (23,24), have been performed. Compaction of long DNA by natural HOs and the behavior of such a biologically relevant system deserves a thorough investigation. The possibility of gene regulation by solely controlling the DNA conformational state (folded of collapsed) remains an open question (25).

The present work was motivated by the importance of DNA length in the DNA compaction process and is focused on unimolecular folding of megabase-long reconstituted chromatin. We report the conformational behavior and compaction of chromatin reconstituted from giant T4 phage DNA (165.6 kb, contour length ca. 60 \(\mu\)m) and recombinant human HOs as a function of the degree of
loading of DNA by HOs and of the nature of polycation added (Mg$^{2+}$ and spermine$^{4+}$), using direct single-molecule DNA visualization in bulk solution by fluorescence microscopy and measurements with the dynamic light scattering method. The reconstituted chromatin in this study, characterized by MNase assay and negative stain TEM, under low salt conditions was shown to form micrometer-sized beads-on-a-string chromatin with a somewhat heterogeneous distribution of nucleosomes on the DNA. The size is close to that of TADs in cell nuclei and this study is the first attempt to address compaction for in vitro reconstituted chromatin fibers of size similar to the TADs of eukaryotic chromatin in vivo. This chromatin reconstituted on long DNA without preferential nucleosome positioning, should serve as a reasonable model of human eukaryotic chromatin because a common characteristic of chromatin in vivo is that the majority of the nucleosomes in the human genome are not preferentially positioned (26). Nucleosomes arrays based on DNA templates with high affinity positioning elements on the other hand, are regularly positioned and are size-limited due to their repetitive design.

**MATERIALS AND METHODS**

The linear T4GT7 DNA was purchased from Nippon Gene Co. Ltd. (Japan). The molar concentration of DNA phosphate groups (i.e. DNA monomer units, nucleotides). The fluorescent dye YOYO-1 (1,1′-(4,4,7,7-tetramethyl-4,7-diazauacene-methylene)-bis-[3-methyl-2,3-dihydro-(benzo-1,3-oxazole)-2-methylidenem]-quinolinium tetraiodide) was provided by Molecular Probes (Invitrogen, Japan). NaCl, MgCl$_2$, and spermine tetrahydrochloride were purchased from Nacalai Tesque Inc. (Kyoto, Japan). Deionized water (Milli-Q, Millipore) was used for samples preparation. TE-KCl buffer (10 mM Tris–HCl, 1 mM EDTA, pH 7.8) containing 10 mM of KCl was used in all experiments, except for transmission electron microscopy where Tris was replaced by HEPES.

**Preparation of recombinant chromatin**

The coding sequences of recombinant human core histones H2A, H2B and H3.1 were cloned in the pET21a vector, and the H4 histone coding sequence in the pET3a vector. Each core histone was individually expressed in the Escherichia coli BL21(DE3)-pLysS strain, purified from inclusion bodies using size-exclusion and cation-exchange chromatography following the established protocol (27).

The HOs were refolded from the equimolar mixture of core histones, purified using size-exclusion chromatography following established protocol (27) and assessed on SDS-PAGE (Supplementary Figure S1). Reconstitution of the T4 DNA with HO was performed using the salt dialysis approach (17,18,28,29). DNA complexes with HO at different degrees of loading were reconstituted by adding 11%, 16%, 23%, 32%, 45%, 66%, 90%, 113%, 128%, 131%, 163% and 200% of HO, calculated based on a ratio of one HO per 200 bp of DNA. For example, 100% loading corresponds to 828 HO molecules per DNA molecule. A typical chromatin reconstitution mixture of 50 μl contained DNA at 0.1 μg/μl, 2 M KCl, 20 mM Tris pH 7.5, 1 mM EDTA pH 8 and 1 mM DTT. The reconstitution mixture was loaded into a Slide-A-Lyzer MINI dialysis unit (3500 MWCO, Thermo Scientific) and dialyzed at room temperature at constant mixing in 200 ml of the initial buffer containing 2 M KCl, 20 mM Tris, 1 mM EDTA and 1 mM DTT.

**UV–Vis spectroscopy**

DNA concentration was measured in samples after chromatin reconstitution with a NanoDrop 2000 UV–Vis Spectrophotometer (Thermo Scientific, NanoDrop Products, Wilmington, Delaware, USA) at 25°C.

**Dynamic light scattering (DLS)**

Dynamic light scattering (DLS) experiments were conducted using a Zetasizer Nano ZS (Malvern Instruments Ltd, Worcestershire, UK) at 25°C. A laser beam was passed through a 1 ml quartz cell containing 0.66 ng/μl (2 μM) T4 DNA in buffer with an appropriate amount of condensation agent (spermine or MgCl$_2$), and the scattered light was measured at a 173° angle to the incident beam. The hydrodynamic size of compact DNA condensates was obtained as a Z-average value (cumulus mean).

**Fluorescence microscopy (FM)**

Sample solutions for single molecule observations were prepared by successive mixing of water, concentrated solution of chromatin at the final DNA concentration of 0.066 ng/μl (0.2 μM), fluorescent dye YOYO (20 nM), and a solution of the multivalent cation as the compaction agent. Solution components were gently mixed after addition of a new component and the final samples were incubated for 1 hour at ambient temperature prior to observations. FM observations were performed using an Eclipse TE2000-U (Nikon, Japan) microscope equipped with 100 × oil-immersion lens. Fluorescent images were recorded and analyzed using an EB-CCD camera and an Argus10 image processor (Hamamatsu Photonics, Japan).

**Transmission electron microscopy (TEM)**

Square mesh copper grids (Motaform 400, EMS) were coated by 10 nm carbon layer using Leica EM ACE600. Chromatin at 5 ng/μl was fixed by 0.01% glutaraldehyde on ice for 10 min, loaded onto the freshly glow-discharged grids, and negatively stained by 2% uranyl acetate. The grids were air-dried at room temperature and observed with a Tecnai T12 microscope equipped with Eagle 4K CCD camera (FEI).

**Atomic force microscopy (AFM)**

A 5 μl droplet of reconstituted T4 chromatin at 32 ng/μl was spotted onto a silica surface. After 10 min to allow for DNA adsorption onto the surface, the specimens were developed by flushing them with ultra-pure water followed by drying in a stream of N$_2$ gas. Atomic force imaging was...
done with a Bruker Dimension Icon microscope (Billerica, MA, USA) at room temperature in air. Images were acquired in the tapping mode with silicon (Si) cantilevers (spring constant of 19–55 N/m) and operated below their resonance frequency (typically 270–370 kHz). The images were flattened, and the contrast and brightness were adjusted for optimal viewing conditions.

**Pulsed-field gel electrophoresis**

The electrophoretic mobility shift of T4 DNA reconstituted with histone octamers was assessed using pulsed-field gel electrophoresis using CHEF-DR II system (BioRad). Equivalent of 0.2 μg DNA with 5% sucrose were loaded into 0.7% Megabase agarose (BioRad) in 0.2× Tris–borate–EDTA buffer and separated at 3 V/cm with switch gradient from 60 to 120 s over 24 h at 14°C.

**Micrococcal nuclease digestion of reconstituted chromatin**

Chromatin was digested by Micrococcal Nuclease (MNase) (New England Biolabs) following established procedure with modifications (30). 16 μg of DNA were digested for 10 min at room temperature following manufacturer’s guidelines using 0.05 U/μl and 0.3 U/μl of MNase. Digestion was stopped by addition of EDTA to 100 mM final concentration. After the addition of 25 μg of Proteinase K (ThermoFisher Scientific), the solution was incubated at 50°C for 30 min. The DNA was purified on a Wizard SV Gel and PCR clean-up system (Promega) and run on 10% polyacrylamide gel electrophoresis at 100 V at room temperature.

**Nucleosome occupancy analysis of T4 DNA sequence**

The sequence analysis was performed using an algorithm available online (31). The sequence of T4GT7 DNA (GenBank: KJ477686.1) was divided into five fragments with overlapping regions of 5250 bp. Default parameters for in vitro reconstitution were used: amplitude 0.2; period 10.1 bp; window 74 bp; and potential −1.0 kT. Regions of DNA sequences with values of nucleosome occupancy higher than 0 indicate areas of high nucleosome affinity with nucleosome positioning effect.

**RESULTS**

**Validation of nucleosome formation on T4 DNA in in vitro reconstitution at different degrees of loading with histone octamer**

In contrast to a vast number of studies of nucleosome arrays that were performed on relatively short DNAs, T4 DNA used in the present work contains 165,660 bp that are sufficient to form ~900 nucleosomes on the DNA chain. A series of DNA complexes with HO in the loading range of 11–200% were reconstituted using the salt dialysis approach, based on a ratio of one histone octamer per 200 bp of DNA. Variation of DNA loading with HO allows for a sampling of different average nucleosome repeat lengths (NRL), which previously was demonstrated to affect the structure of chromatin fibers on the 10–30 nm level (9,32).

![Figure 2.](https://example.com/figure2.png) In vitro reconstituted T4 DNA with HO at different degree of loading was digested by micrococcal nuclease (lanes 3–10). In lanes 3 and 8–10, low concentrations (0.05 U/μl) of MNase were used (L); in lanes 4–7, high concentrations of MNase (0.3 U/μl) of MNase were used (H). T4 DNA in the absence of HO shows digestion dependence on MNase concentration (lanes 3 and 4). 100 bp marker DNA (lane 2) allows to estimate that DNA in multiples of around 150 bp is protected from MNase digestion (lane 10). Nucleosome array with 601 nucleosome positioning sequence of 36 nucleosomes with 177 bp nucleosome repeat length digested by MNase (lane 1) produces nucleosome ladder upon incomplete digestion and was used as a positive control.

A gradual decrease in the electrophoretic mobility of the reconstituted chromatin was observed using pulsed-field gel electrophoresis (PFGE) indicating the increased binding of HO to DNA with higher HO loading (Supplementary Figure S2).

Micrococcal nuclease that digests DNA not bound by proteins (33) was used to demonstrate the formation of nucleosomes. Distinct DNA protection band pattern of around 150 bp after MNase digestion results from DNA wrapped around the HO (Figure 2). MNase digestion is highly sensitive to the conditions of the reaction, such as the amounts and molecular weight of DNA, amount of MNase, duration and temperature. Two concentrations of MNase that allow better characterization of reconstituted chromatin at different loading degrees of HO were used. Using high concentration of MNase results in sufficient digestion of DNA and visualization of the 150 bp DNA protection pattern at both low and high loading degrees of HO in the range of 16–131% (lanes 4–7 on Figure 2, see also Supplementary Figure S3A). At low concentration of MNase, partial digestion of DNA and chromatin at low degrees of HO loading results in smeared higher molecular weight DNA bands (Supplementary Figure S3B). At HO loading degrees of 90% and 113% around the saturation, partial digestion results in a ladder pattern in multiples of 150 bp, indicating the release of not only mononucleosomes, but also di-, tri-, and tetranucleosomes (lanes 8 and 9 on Figure 2, see also Supplementary Figure S3B). Because the T4 DNA is not a positioning sequence, unlike the control nucleosome array with the 601 sequence of 36 nucleosomes with 177 bp NRL (Figure 2, lane 1), the difference in MNase digestion is not unexpected. Partial MNase digestion, accompanied...
by DNA smears that allow detection of tetraneucleosomes as the largest product that can be resolved, indicates a heterogeneous distribution of the NRL in T4 chromatin (see below on TEM results). This is in agreement with observations for MNase digestion of nucleosome arrays reconstituted on telomeric DNA formed by repetitive TTAGGG repeats that lack positioning elements (34). At higher saturation ratios, digestion products were not observed. Hereafter, we reserve the term ‘saturated chromatin’ for those with saturation degree in the 90–128% range, ‘subsaturated chromatin’ for those with lower HO contents, and ‘oversaturated’ for higher HO loading values.

Transmission electron microscopy (TEM) and atomic force microscopy (AFM) techniques were used to confirm the formation of multiple nucleosomes on T4 DNA. Chromatin at 100% of HO loading under low salt conditions was observed as a coil-like structure (35) (Figure 3). The resolution on the TEM image (Figure 3A) allows two turns of DNA wrapped around the HO to be distinguished for the individual nucleosomes in a side view orientation. Individual nucleosomes of ca. 10 nm size (region 1) connected by a linker DNA of variable length appearing as beads-on-a-string (region 2), and as aggregated and stacked nucleosomes (region 3) were also identified on the TEM image (Figure 3A). What appears as a single chromatin molecule imaged by AFM displaying a beads-on-a-string 10 nm fiber conformation, exhibits heterogeneous distribution of nucleosomes along the DNA chain (Figure 3B). The imaged structure of recombinant T4 chromatin in this work is in agreement with the beads-on-a-string structure of 100 kb DNA reconstituted with purified HOs and imaged using AFM (17,18) and the heterogeneous structure of rat liver chromatin visualized using TEM from a recent study (36).

The T4 DNA sequence that may influence nucleosome distribution was analysed using a program optimized for in vitro reconstitution (31). The probability of nucleosome occupancy over the whole length of T4 DNA was observed to have a mean value of −1.8 with standard deviation ± 0.64 (Supplementary Figure S4). Occupancy values higher than 0, indicative of high affinity for nucleosome formation according to the authors’ model, were not detected. The observed occupancy values in the negative range may be interpreted as sequences with nucleosome formation ability without positioning preference, which would allow random distribution of nucleosomes on the DNA for subsaturated systems. The DNA sequence should therefore not have a negative effect on nucleosome formation, because nucleosome free regions have been shown to form as a result of the action of chromatin remodelers, and not due to the DNA sequence (37). The observed nucleosome occupancy of T4 DNA, therefore leads to formation of nucleosome arrays where the degree of loading is likely to be related to the average nucleosome repeat length.

**Conformational behavior of single T4 DNA molecules reconstituted with histone octamers**

The long contour length of T4 DNA (ca. 60 μm) makes it possible to visualize reconstituted complexes at the level of individual molecules by fluorescence microscopy (FM) techniques. T4 DNA and chromatin complexes were labeled by the fluorescent dye YOYO and their conformational behavior was monitored in bulk solution by FM. Figure 4A shows typical snapshots of a single DNA molecule in a coil conformation exhibiting Brownian motion in a bulk solution of TE buffer with 10 mM KCl. The measured DNA long-axis length (L), which is defined as the longest distance in the outline of the DNA fluorescent image, is larger than the actual size by ~0.3 μm in each dimension (Figure 4B) due to a ‘blurring effect’ attributed to the resolution limit of FM (38). Upon reconstitution with HOs, the DNA coil gradually shrank (Figure 4A and C) from ca. 3.5 to 2 μm but preserved Brownian motion as well as intramolecular fluctuations.

The shrinking of DNA occurs due to the wrapping of DNA around HOs according to the well-established structure implying wrapping of ca. 50 nm of DNA length per one HO of ca. 7 nm diameter. The average values of the DNA long-axis length are plotted as a function of a loading degree in Figure 4D. As a result of nucleosome formation, the long-axis length of DNA decreased ~1.7-fold, which corresponds to about a 5-fold decrease of the DNA coil molecular volume. Based on FM observations, single molecules of DNA reconstituted with HOs are observed up to 128% loading. Further increase in HO loading induces intermolecular aggregation (data not shown).

**Compaction of reconstituted chromatin by divalent cations**

The conformation of the nucleosome arrays is strongly affected by the presence of mono- and multivalent cations (5) which have been used in studies to induce aggregation and precipitation of nucleosomes and chromatin (39–43). Therefore, the conformational behavior of chromatin reconstituted from T4 DNA in the presence of Mg2+ ions was studied. Figure 5 shows the dependence of the long-axis length of DNA and reconstituted chromatin on the concentration of Mg2+ ions in solution. According to Manning counterions condensation theory (44) that was applied to DNA in solution by Bloomfield and colleagues (45), divalent cations are unable to compact DNA because DNA anionic charges are not sufficiently neutralized by the relatively weakly bound divalent counterions. In good agreement with condensation theory and earlier experimental studies, the coil conformation of T4 DNA was preserved and the long-axis length of DNA only slightly decreased in solution with Mg2+ concentrations of up to 10 mM. Due to the competition of Mg2+ and the YOYO fluorescent dye for DNA binding, higher Mg2+ concentrations were not studied by FM.

The dynamic light scattering (DLS) technique was used to examine effects of higher Mg2+ concentrations and to gain additional information about T4 DNA phase transition at the level of the molecular ensemble. DLS results in Figure 6 indicate that no DNA phase transition occurred up to 100 mM Mg2+. The conformational response of chromatin due to Mg2+ addition at 23% of HO loading observed by FM was similar to that of free T4 DNA (Figure 5). The DLS measurements, however, revealed an increase in scattered light intensity for chromatin at 11% and 23% of HO loading above ca. 30 mM of Mg2+ that may be attributed to either intermolecular aggregation or intramolecular con-
Figure 3. *In vitro* reconstituted T4 chromatin at 100% HO loading under low salt conditions were imaged as coil-like structures using TEM (A) and AFM (B). Regions 1, 2 and 3 indicate a single nucleosome, beads-on-a-string, and aggregated nucleosomes, respectively. Scale bars are 300 nm (A) and 500 nm (B).

Figure 4. Single-molecule FM observations of T4 DNA conformational changes upon reconstitution with HOes. (A) Series of typical fluorescence micrographs of a YOYO-labeled single T4 DNA molecules and T4 DNA reconstituted with HOes at loading degrees 45% and 128% observed in a bulk solution of TE buffer with 10 mM of KCl. The time interval between snapshots is ca. 0.5 s. (B) Schematic representation of the relationship between actual DNA molecule in a coil or in a globule conformation and the correspondent fluorescent images, that are larger than actual DNA chain by ca. 0.3 μm due to blurring effect, and the definition of the characteristic parameter of long-axis length (L). (C) Long-axis length distributions of T4 DNA molecules and T4 DNA reconstituted with HOes at loading degrees 45% and 128% under conditions of A. (D) Changes in the average long-axis length of T4 DNA and DNA reconstituted with HOes as a function of a loading degree. Experimental points of different color correspond to two independently prepared samples. The error bars indicate the standard deviations of the average values measured over at least 100 individual DNA or chromatin molecules.
Compaction of T4 DNA and reconstituted chromatin by divalent cation (Mg$^{2+}$). Changes in the average long-axis length of YOYO-labeled T4 DNA (0.2 $\mu$M) and T4 DNA reconstituted with HO$\theta$s (0.2 $\mu$M) at different loading degrees at various concentrations of magnesium chloride (MgCl$_2$) in a bulk solution of TE buffer with 10 mM of KCl. Blue areas correspond to Mg$^{2+}$ concentrations at which DNA or chromatin molecules are completely compacted (globule state). The error bars indicate the standard deviations of the average values measured over at least 100 individual DNA or chromatin molecules.

For HO loading degrees of 45% and higher, the conformational change from the unfolded coil to the compact globule and the increase in scattered light intensity were observed by FM and DLS, respectively (Figures 5 and 6). While DLS profiles for these chromatin complexes are very similar (Figure 6), single molecule FM analysis (Figure 5) shows complete compaction into globules (shaded areas) taking place at a lower concentration of Mg$^{2+}$ ions for chromatin with higher HO loading, reaching ca. 2 mM value for the most saturated complex at 128%. The important characteristic of chromatin compaction upon addition of Mg$^{2+}$ ion was the constant long-axis length of the chromatin coil before the onset of compaction at around 1 mM of Mg$^{2+}$ in all samples. It should be noted, that due to a large number of HO bound to individual T4 DNA molecules, a certain heterogeneity in chromatin composition can result in a slight variation of the conformational transition point or compaction mechanism for individual chromatin molecules.

Hydrodynamic size of compacted chromatin molecules in the 90–128% HO loading range was measured by the DLS (Supplementary Figure S5A). The size of saturated chromatin remains constant, ca. 250 nm, in a rather broad concentration range of 4–60 mM Mg$^{2+}$, indicating the intramolecular compaction being the predominant mechanism responsible for the observed increase in scattered light intensity.

**Spermine is unable to induce a first-order phase transition of chromatin**

Next, we studied compaction of DNA and reconstituted chromatin by the tetravalent cation, spermine. Bearing a charge higher than +2, spermine is able to induce compaction of free DNA by providing a sufficient charge neutralization of the DNA polyanion (45). The DLS measurements showed that T4 DNA and chromatin molecules of all studied saturation degrees underwent compaction at very similar concentrations of spermine (Supplementary Figure S6). However, the DLS method is unable to account for a far more intriguing characteristic of DNA compaction—the DNA folding scenario at the level of single DNA molecules. The striking feature of DNA compaction by tetravalent cations, the discrete conformational change at the level of single molecules, has been reported more than two decades ago (20), and in this study the mechanism of chromatin compaction was closely examined.

Figure 7 shows compaction curves for T4 DNA and chromatin of different HO loading degrees. Increase of the HO loading degree has a negligible effect on the chromatin compaction efficiency of spermine. Complete compaction of DNA and chromatin into globules of ca. 0.5 $\mu$m takes place at ca. 20 $\mu$M of spermine. However, the compaction mechanism for T4 DNA and chromatin at 11% HO loading was drastically different from the compaction mechanism of chromatin at loading degrees of 23% and higher. While the compaction for both T4 DNA and chromatin at 11% HO loading occurs as an all-or-none mechanism, at 23% and higher chromatin displays a continuous character of the transition.

Figures 8A and B show representative single-molecule fluorescent images of chromatin at 11% and 23% HO loading and contrast the change of DNA long-axis length distributions at different concentrations of added spermine. Compaction of chromatin at 11% takes place between 10
and 30 μM of spermine as a first order phase transition of DNA conformation from 3.5 μm coil to 0.7 μm globule through a region with a bimodal size distribution (yellow areas on Figure 7) where coil and globule conformations coexist. Coexistence of unfolded and compact forms of DNA preceding DNA collapse is a typical intermediate state in coil-globule transition of a semi-flexible long polyelectrolytes such as DNA and has been repeatedly demonstrated experimentally (46,47) and by computer simulations (48). The coexistence state appears due to double minima in the free energy profile of the DNA conformation. In contrast, the chromatin coil at 23% HO loading undergoes gradual shrinking toward compaction into a 0.7 μm globular condensate (Figure 8). The increase in the HO loading enables chromatin to respond to the addition of small amounts of spermine by gradual compaction.

Although compaction of chromatin by spermine is described as a gradual shrinking, we noticed a large population of segregated structures in the course of the compaction of chromatin complexes at loading degrees of 90% and higher (Figure 9). Segregated structures containing both coil and folded globular parts in the same DNA molecule have been reported earlier for hydrophobic binders (49,50) and interpreted as the discrete coil-globule transition occurred within the same DNA molecule. A large fraction of molecules with intra-chain segregation observed in our system may indicate a heterogeneous HO positioning along DNA that can cause preferential condensation of a DNA part that is more populated by HOs. Contrary to spermine, structures with intra-chain segregation were not formed during the chromatin compaction by Mg²⁺ cations, instead, the coil of DNA shrank gradually and homogeneously to a final globular compact state.

The hydrodynamic size of the compacted chromatin complexes was measured by the DLS technique. Unlike chromatin complexes compacted by Mg²⁺, with constant hydrodynamic size, the size of chromatin condensates compacted by spermine depends on both the degree of DNA loading with HO and the concentration of spermine (Supplementary Figure S5B). Compaction of free T4 DNA yields very compact condensates of ca. 100 nm hydrodynamic diameter in good agreement with earlier studies (51). The increase of spermine concentration induced a slight increase of condensate size to ca. 150 nm. It should be noted that the size of DNA condensate compacted by spermine is significantly smaller than the size of condensates compacted by Mg²⁺.
Figure 7. Compaction of T4 DNA and chromatin by tetravalent cations (spermine). Changes in the average long-axis length of YOYO-labeled T4 DNA and T4 DNA reconstituted with HOs at different loading degrees at various concentrations of tetravalent cation, spermine (Spm4+), in a bulk solution of TE buffer with 10 mM KCl. Experimental points show separately the average sizes of coils and globules for T4 DNA and chromatin at 11% HO loading at intermediate spermine concentrations characterized by a bimodal distribution of DNA long-axis lengths. Yellow and blue areas are regions of coil-globule coexistence and complete DNA compaction, respectively. The error bars indicate the standard deviations of the average values measured over at least 100 individual DNA or chromatin molecules. The percentage values above experimental points in the coexistence region indicate the percentage of T4 DNA or reconstituted chromatin in the compact state.

DISCUSSION

The higher-order structure of chromatin and its manner of compaction are closely associated with its genetic activity in cells (52–54). However, while our present understanding of chromatin condensation/compaction mainly comes from in vitro studies on well-defined systems based on short nucleosomal arrays with nucleosome positioning sequences (55), the chromatin in vivo contains significantly longer DNA chains (up to hundreds of thousands bp in a single chromosome loop) (56) folded hierarchically into heteromorphic structures (57), with the majority of nucleosomes positioned irregularly (26,36,58,59). Therefore, studies of chromatin reconstituted on large DNA molecules without nucleosome positioning will improve our understanding of chromatin properties in vivo.

FM observations show that 165.6 kb giant DNA folding into chromatin is accompanied by a gradual shrink-
Figure 8. Two compaction scenarios of T4 chromatin by a tetracation. (A) Typical fluorescence micrographs of a YOYO-labelled T4 DNA reconstituted with HOs at 11% (left) and 23% (right) loading degrees, observed at different concentrations of spermine in a bulk solution of TE buffer with 10 mM of KCl. The concentrations of spermine are indicated above corresponding images. (B) Long-axis length distributions of chromatin at 11% (left) and 23% (right) loading degrees under conditions of A.

Figure 9. Formation of intrachain segregation structures during compaction of reconstituted chromatin. Series of typical fluorescence micrographs of a YOYO-labelled chromatin at 113% loading degree in the presence of 6.7 μM spermine in a bulk solution of TE buffer with 10 mM of KCl. The time interval between snapshots is ca. 0.5 sec.
ing of the DNA coil dependent on the HO loading (Figure 4). At the level of the resolution of the FM experiment, the fluorescence intensity was homogeneously distributed along individual coils of both subsaturated and saturated chromatin molecules, indicating no strong heterogeneity of nucleosome density along DNA chains and the sequence-based probability of nucleosome occupancy (Supplementary Figure S4). However, the HO positioning on the DNA chain was characterized by a certain heterogeneity manifested in a difference in linker DNA length (Figure 3). Over-saturation of DNA with HOs induces intermolecular aggregation, likely involving nonspecific binding of HOs to DNA and nucleosomes.

The bioavailable polycations Mg\(^{2+}\) and spermine exist at millimolar concentrations in the living cell (60,61), thus they can be directly involved in regulation of chromatin higher-order structure in vivo. Although one may expect similar trends in compaction of negatively charged chromatin by divalent and tetravalent cations as a result of neutralization of negatively charged DNA phosphates, our results reveal a considerable difference between the effects of divalent magnesium cations and tetravalent spermine cations on the higher-order structure of subsaturated and saturated nucleosomal arrays, in agreement with cation-dependent effects reported earlier (43).

The protection from MNase digestion indicates the compacted state of chromatin. The observed protection of chromatin may result from the close positioning of nucleosomes on DNA, and nonspecific binding of histones. Furthermore, the presence of 5 mM Ca\(^{2+}\) cations in the MNase reaction buffer augments compaction and aggregation. Compacted chromatin globules were observed in the millimolar range of Mg\(^{2+}\) concentrations by FM in this work (Figure 5). For oversaturated chromatin, at ratios higher than 131\% the excess cationic HOs may bind non-specifically to the negatively charged chromatin fibers, leading to chromatin neutralization and aggregation. At degrees of HO loading above 128\%, the chromatin may represent a beads-on-a-string structure with very close-packed nucleosomes, which may not be the most physiologically relevant complex. Although telomeric chromatin has been shown to have 157 nucleosome repeat length (62), the majority of eukaryotic chromatin has a nucleosome every 200 ± 40 bp (63).

**Chromatin compaction by magnesium ions**

Mg\(^{2+}\) ions do not induce condensation of double-stranded DNA because its electrostatic interaction with DNA does not provide sufficient DNA charge neutralization (64). On the other hand, Mg\(^{2+}\) is known to effectively promote condensation/aggregation of nucleosomes and nucleosome arrays (42,65). Thus, Mg\(^{2+}\)-induced aggregation of nucleosomes, in which the DNA's charge is partly neutralized by cationic histone octamers (40), is due to attractive interactions. Attractive interactions between nucleosomes in chromatin originate from the ion-ion correlation effects (66) due to a positional correlation of Mg\(^{2+}\) ions condensed on different adjacent negatively charged segments of DNA (67) in nucleosomes, coupled with histone tail bridging (5). A certain contribution to minimize the free energy in the chromatin compaction process originates from an increase of translational entropy of counterions (68) released during chromatin compaction and effects associated with cationic tails (69).

Because the saturated chromatin can be compacted by Mg\(^{2+}\) in contrast to 'naked' DNA (Figure 5), the degree of chromatin saturation by histones is clearly the factor that determines the ability of a divalent cation to compact chromatin. Indeed, in earlier systematic studies on subsaturated nucleosome arrays, the degree of saturation was shown to play a crucial role in the chromatin condensation (70). The condensation of subsaturated chromatin by Mg\(^{2+}\) was significant only if saturation degrees were higher than ca. 70\% (70,71). This saturation degree threshold was explained based on pairwise preference in nucleosome occupation (72). These studies emphasized the importance of adjacent nucleosomes interaction for chromatin condensation. The saturation degrees above ca. 70\% imply the absence of large gaps between adjacent nucleosomes providing effective inter-nucleosomal interactions through histone cationic tails leading to the efficient chromatin condensation.

The long chromatin compaction in our study has many similarities with earlier results on well-defined short nucleosome arrays. Free DNA is not compacted by divalent cations (Figures 5 and 6). Due to the relatively weak interaction of Mg\(^{2+}\) ions with both DNA and nucleosomes, the conformation of chromatin remains virtually unchanged until millimolar amounts of Mg\(^{2+}\) are added to induce association of the nucleosomes. In contrast to short fragments of subsaturated chromatin (70,71), Mg\(^{2+}\) induced condensation of highly unsaturated chromatin at 11\% of HO loading was detected by DLS technique (Figure 6), but at Mg\(^{2+}\) concentrations 10 times higher than sufficient to induce the compaction of the saturated chromatin at 113\% and 128\%. The concentration of Mg\(^{2+}\) needed to induce single-molecule compaction of chromatin with loading higher than 40\% decreased moderately and approached concentration of several mM – typical Mg\(^{2+}\) concentration to induce association of saturated nucleosome arrays (73,74) or aggregation of nucleosomes in solution (65). Despite the very low concentrations used here for megabase-long chromatin in the FM observations, the compaction behavior is similar to the intermolecular self-association (aggregation) observed for reconstituted 12-mer arrays using the ‘601’ positioning sequence (67). The onset and the cooperativity of nucleosome array aggregation and the compaction of single chromatin molecules of high saturation degree occur at comparable Mg\(^{2+}\) concentrations. Condensation of chromatin begins in both cases at ca. 1 mM of Mg\(^{2+}\) added to the solution and ended at 4 mM Mg\(^{2+}\) for 12-mer nucleosomal array (74) and at 4–7 mM for saturated T4 chromatin (Figures 5 and 6).

The results above clearly show that the high degree of chromatin saturation is the essential condition for efficient chromatin compaction by Mg\(^{2+}\) regardless of chromatin length. Nevertheless, compaction of long chromatin by Mg\(^{2+}\) can be achieved even at low degrees of saturation, at which short nucleosome arrays associate poorly (75). This observation may be relevant for chromatin in the cell, where chromatin containing several hundreds of nucleosomes may have the histone-depleted DNAs regions...
in a compacted state because of a sufficient number of nucleosome-nucleosome attractive interactions.

In a recent paper, Maeshima et al. investigated condensation of short nucleosomal arrays by Mg\(^{2+}\) cations and found that a great number of molecules cooperatively assemble into globular condensates with sizes between 50 and 1000 nm (76). In contrast, under the present conditions, compaction of megabase-long chromatin, induced by Mg\(^{2+}\) occurs as a single molecule event resulting in formation of globules of ca. 250 nm with no detectable aggregation tendency into larger condensates. The main reason is that under the present conditions of very low DNA concentration, monomolecular collapse of chromatin is thermodynamically preferred to intermolecular aggregation (77–79).

Despite the general agreement in concentration ranges of Mg\(^{2+}\) and sizes of aggregates observed in both studies, the long DNA used in our work may be more biologically relevant, thus being a suitable in vitro model system to study chromosome structure and functions.

Previous studies of native chromatin containing ~40 nucleosomes per DNA chain also acknowledged the formation of 30 nm chromatin fibers (80) preceding compaction of chromatin into globules, yet the relevance of 30 nm fiber to chromosome assembly in vivo is recently under debate in the field (81). Chromatin compaction profiles (Figure 5) and DLS data (Figure 6) show the saturated chromatin coil undergoing a gradual decrease in size while it turns into a globule. DLS measurements of the hydrodynamic radii (Supplementary Figure S5A) revealed only one type of compact condensate of several hundred nanometers in size, which corresponds to a globular form of condensed chromatin. All these observations support a model of direct folding of dynamic 10-nm chromatin fiber into a globular condensate although the formation of regions of 30 nm fiber may not be ruled out.

Chromatin compaction by spermine

Compaction of chromatin by tetravalent cations is drastically different from Mg\(^{2+}\)-induced compaction. Considering the purely electrostatic effect of spermine on nucleosome aggregation, it is clear that spermine induces nucleosome aggregations at lower concentrations than Mg\(^{2+}\). According to Manning’s theory (44), the fraction of neutralized charges on free DNA is equal to 94% in the presence of tetravalent cations, which is sufficient to induce DNA compaction that requires ca. 91% charge neutralization (64).

The same theory gives 96% of phosphates neutralized by spermine for DNA in nucleosomes (40). Consequently, not only spermine can induce aggregation of nucleosomes as we discussed for Mg\(^{2+}\)-induced chromatin compaction, but its interaction with DNA can also largely promote attractive interactions between segments of free DNA. Because the neutralization of free DNA charges and DNA in nucleosomes by spermine is of the same degree, free DNA and chromatin of different saturation degree should fold at approximately the same amount of added spermine to solution. Both DLS and FM studies show constant spermine concentration (ca. 20 \(\mu\)M) at which DNA and chromatin are folded into compact globules (Figure 7 and Supplementary Figure S6).

There is, however, a fundamental difference between the compaction mechanisms at the level of single molecules. The free DNA undergoes a discrete coil-globule transition (20), whereas the saturated chromatin is observed to shrink gradually (Figures 8 and 9). For subsaturated chromatin samples, the change of compaction mechanism from discrete to continuous occurs between 11% and 23% of HO loading (Figure 7).

Earlier studies have shown that when a polymer chain is rigid enough, its folding transition occurs in a discrete manner (82), whereas the transition is continuous for flexible polymeric chains. Free DNA with a persistence length of 50 nm (83) is classified as a relatively rigid (or semi-flexible) polymer chain, while the nucleosomal arrays and chromatin are more flexible with a persistence length <30 nm according to single-molecule experiments (84,85). Therefore, the alteration of the effective persistence length of DNA by the degree of histone saturation is expected to affect the manner of chromatin folding by multivalent cations.

The discrete transition of the DNA coil to a globule results in a very tight packaging of DNA similar to the DNA packaging in bacteriophage heads (90) and causes a complete inhibition of DNA transcriptional activity (91). In this regard, DNA binding with only 20% of HO compared to the saturation amount leads to a change in DNA compaction scenario that may allow for a more flexible regulation of DNA genetic activity in eukaryotic cells.

Another important feature of chromatin compaction by spermine revealed in our FM study is a strong intra-chain segregation along individual chromatin molecules with a high degree of nucleosome saturation (92). In aggregation studies of short chromatin fragments, it was demonstrated that DNA saturation by HO is a stochastic process at low and medium saturation degrees, while a certain weak cooperativity is manifested at high saturation degrees due to pairwise preference in nucleosome occupation (72) caused by histone tail interactions (92). Atomic force microscopy studies by Yoshikawa et al. also revealed a bimodal profile of nucleosome densities along reconstituted chromatin of high saturation degrees (52), that supports the mechanism of a cooperative HO binding when the density of HO is
high. Therefore, one may expect that chromatin at HO loading 90% and higher possesses a certain degree of heterogeneity in terms of nucleosome density. This heterogeneity leads to a fluctuation in a charge density and, probably, structure of chromatin along the same DNA that causes a difference in chromatin compaction states at the same concentration of multivalent cation. In good agreement with this model, strong intra-chain segregation was observed only for chromatin with high degree of saturation (90% and higher) while no intra-chain segregation was observed for subsaturated chromatin. This phenomenon might have an important implication to gene regulation: cooperative nucleosome positioning may result in more efficient compaction of such chromosomal regions resulting in suppression of genes over a significant DNA length. The observation of intrachain segregation or nucleosome clustering observed in our re-combinant system by FM and AFM, may mimic the organization of chromatin in situ (59) and ‘nucleosome clutching’ (58) reported recently. A thorough characterization of our model system is needed before a mechanism responsible for nucleosome segregation and its relevance to the in vivo chromatin structure and function can be established.

We noted that intra-chain segregation was observed only in chromatin compaction by spermine, but not by Mg$^{2+}$. This difference can be explained by insufficient stabilization of a compact chromatin phase due to weak condensation potency of Mg$^{2+}$ in comparison with spermine. Spermine-mediated compaction, in turn, better mimics chromatin compaction in nuclei by polycondensed, like histone tails, and linker histones.

CONCLUSIONS

By single-molecule observations of DNA and chromatin interacting with multivalent cations, we uncovered hitherto unknown features of unimolecular DNA compaction. The most important findings are the following: (i) Complexation of DNA with a small fraction (10–20%) of histones is sufficient to induce DNA compaction by adding divalent cations such as Mg$^{2+}$. (ii) Complexation of DNA with a small fraction (ca. 20%) of histones is sufficient to change the first-order DNA compaction transition into a gradual shrinking to a compact globule. (iii) Compaction of saturated chromatin by spermine proceeds through intrachain segregation in single DNA molecules containing unfolded and compact parts in the same chain. This first study of reconstituted chromatin of size similar to TADs, i.e. nearly megabase length of DNA and ca. 300 nm size of chromatin condensate, is a step forward to reproducing the state of chromatin in the nucleus and hence of great biological relevance as a model for chromatin condensation in vivo.

Studies on genome-wide nucleosome positioning revealed that most of eukaryotic chromatin is heterogeneous and does not show regular nucleosome positioning on the same DNA sequences in different cell types or during cell development stages or on functionally similar DNA sequences (26). Therefore, the somewhat heterogeneous chromatin reconstituted on a long nonspecific DNA template like T4 DNA in the present study, should be a more relevant in vivo model for chromatin formed on large megabase size DNA, compared to saturated nucleosomal arrays with regular NRL, constructed using DNA template based on high affinity nucleosome positioning sequences, which are limited in size.

SUPPLEMENTARY DATA

Supplementary Data are available at NAR Online.

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